





## Draft Genome Sequences of Lactobacillus salivarius A3iob and Lactobacillus johnsonii CRL1647, Novel Potential Probiotic Strains for Honeybees (Apis mellifera L.)

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ABSTRACT This report describes the draft genome sequences of Lactobacillus salivarius A3iob and Lactobacillus johnsonii CRL1647, probiotic strains isolated from the gut of honeybee Apis mellifera workers. The reads were generated by a wholegenome sequencing (WGS) strategy on an Illumina MiSeq sequencer and were assembled into contigs with total sizes of 2,054,490 and 2,137,413 bp for the A3iob and CRL1647 strains, respectively. The draft genome sequences of L. salivarius A3iob and L. johnsonii CRL1647 will be useful for further studies of the specific genetic features of these strains and for understanding the mechanisms of their probiotic properties.

actobacillus salivarius subsp. salivarius A3iob and Lactobacillus johnsonii CRL1647 were isolated from the gut of honeybee Apis mellifera workers and selected among several isolates due to their antimicrobial properties and high lactic acid production (1, 2). Both Lactobacillus strains elicited probiotic properties when administered individually to commercial productive hives of the honeybee Apis mellifera L. Lactobacillus strains generated a higher number of bees through stimulation of the queen's egg laying (1-3). An increase in the resistance to parasitic mite (Varroa spp.) and microsporidian parasite (Nosema spp.) infections has been also observed after the administration of A3iob or CRL1647 strains (1, 4). These beneficial properties have been evaluated and confirmed in different ecoregions of Argentina and under several different beekeeping practices (5).

L. salivarius A3iob and L. johnsonii CRL1647 were cultured for 12 h at 37°C (final log phase) in de Man-Rogosa-Sharpe broth (MRS; Oxoid, Cambridge, UK), and genomic DNA isolation was performed as described by Azcárate-Peril and Raya (6). Draft genome sequences of both bacteria were obtained with an Illumina MiSeq platform using the  $2 \times 150$ -bp paired-end read length sequencing protocol. The *L salivarius* A3iob and *L*. johnsonii CRL1647 data sets contain 2,216,287 and 2,391,328 reads, respectively. The raw sequence data were analyzed by FastQ for quality control purposes. These reads were de novo assembled with SPAdes version 3.11.1 (7). The A3iob strain contained 12 contigs (2,054,490 bp, 34.6% G+C content,  $114.0 \times$  coverage), while the CRL1647 strain contained 38 contigs (2,137,413 bp, 34.4% G+C content,  $72.0\times$  coverage). The Rapid Annotations using Subsystems Technology (RAST) server and standalone Prokka (rapid prokaryotic genome annotation) program were used for functional annotation of predicted genes (8, 9). A total of 1,911 coding sequences, 61 tRNAs, 20 rRNAs, 3 noncoding RNAs (ncRNAs), and 1 CRISPR array were annotated in the L salivarius A3iob genome.

Received 10 July 2018 Accepted 24 July 2018 **Published** 16 August 2018

Citation Audisio MC, Albarracín L, Torres MJ. Saavedra L, Hebert EM, Villena J. 2018. Draft genome seguences of Lactobacillus salivarius A3iob and Lactobacillus johnsonii CRL1647, novel potential probiotic strains for honeybees (Apis mellifera L.). Microbiol Resour Announc 7:e00975-18. https://doi.org/10.1128/MRA

Editor Frank J. Stewart, Georgia Institute of

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In the *L. johnsonii* CRL1647 genome, 1,762 coding sequences, 30 tRNAs, 6 rRNAs, 3 ncRNAs, and 2 CRISPR arrays were found.

The genomes were further analyzed with BAGEL4 for the detection of bacteriocin genes (10). Salivaricin and enterolysin A genes were found in the A3iob genome, while helveticin J, thermophilin A, and enterolysin A genes are present in the CRL1647 genome. Both *Lactobacillus* strains contain genes encoding fibronectin binding protein that could be involved in their probiotic effect. In addition, clusters of genes involved in the biosynthesis of pyridoxine, folate, biotin, and riboflavin were found in the A3iob genome, while the CRL1647 genome harbors genes involved in folate, thiamine, and riboflavin biosynthesis.

The draft genome sequences of the A3iob and CRL1647 strains will be useful for further studies of their specific genetic features and for understanding the mechanisms of their probiotic properties.

**Data availability.** The draft genome sequences of *Lactobacillus salivarius* subsp. *salivarius* A3iob and *Lactobacillus johnsonii* CRL1647 have been deposited as wholegenome shotgun sequencing projects at DDBJ/EMBL/GenBank under the accession numbers QFAS00000000 and QFBA00000000, respectively. The versions described in this paper are the first versions, QFAS01000000 and QFBA01000000.

## **ACKNOWLEDGMENTS**

This study was supported by ANPCyT-FONCyT grants PICT-2013-3219 and PICT-201-0410 to Julio Villena, and grant PICT-640-2015 to Marcela Carina Audisio.

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